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APPENDIX III

**COPY OF PROTOCOL** 

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HTR Study No.: 02-120918-106 Kimberly-Clark Corporation

HILL TOP RESEARCH, INC.

HILL TOP RESEARCH CONFIDENTIAL

## PROTOCOL FOR

MODIFIED AOAC GERMICIDAL AND DETERGENT SANITIZING ACTION OF DISINFECTANTS – Rough Plastic One Step Cleaner Sanitizer

For: Kimberly-Clark Corporation

HTR Ref.: 02-120918-106



HTR Ref No.: 02-120918-106 Modified AOAC Germicidal and Detergent Sanitizing Action of Disinfectants Protocol – Rough Plastic Hill Top Research Confidential

### TABLE OF CONTENTS

1.0	INTRODUCTION	.1
2.0	PURPOSE	.1
3.0	STUDY SPONSOR AND SPONSOR REPRESENTATIVE	. 1
4.0	TEST FACILITY AND INVESTIGATIVE PERSONNEL	1
5.0	APPLICABLE REGULATION	2
6.0	RESEARCH STANDARDS	2
7.0	EXPERIMENTAL DESIGN	2
8.0	PROPOSED EXPERIMENTAL STARTING AND EXPERIMENTAL	
	TERMINATION DATES	3
9.0	TEST SUBSTANCE IDENTIFICATION	3
10.0	TEST SUBSTANCE CHARACTERIZATION	3
11.0	TEST SYSTEM JUSTIFICATION	3
12.0	TEST SYSTEM IDENTIFICATION	3
13.0	TEST PROCEDURE	3
14.0	STATISTICAL METHOD	6
15.0	REPORT	6
16.0	DATA RETENTION	6
17.0	NOTICE	7
18.0	PROTOCOL APPROVAL FORM	8
Anne	ndix I/AOAC Method	



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants Protocol - Rough Plastic

Hill Top Research Confidential

### 1.0 INTRODUCTION

Single-use wipes containing chemical sanitizers suitable for use on lightly soiled, nonporous, food contact surfaces are generally tested by a time kill method where the cidal effect of a specific concentration of chemical agent is measured against both a Gram negative and a Gram positive bacterium over a specified time period. The percent reduction in numbers of test bacteria containing a soil load is calculated as compared to a positive control. Standard practices for testing use the AOAC Germicidal and Detergent Sanitizing Action of Disinfectants Method as described in Chapter 6, Disinfectants, Official Methods of Analysis for AOAC International, 17<sup>th</sup> Edition, 2000, Section 6.3.03. This method will be modified to test a single-use wipe on a moderately soiled, rough plastic surface in accordance with the EPA Interim Guidance "Non-Residual Sanitization of Hard Inanimate Food Contact Surfaces Using Pre-Saturated Towelettes (April 12, 2001)".

### 2.0 PURPOSE

To determine the sanitizing action of a wipe containing a chemical agent that can be permitted for use in sanitizing moderately soiled, rough plastic surfaces.

# 3.0 STUDY SPONSOR AND SPONSOR REPRESENTATIVE

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## 4.0 TEST FACILITY AND INVESTIGATIVE PERSONNEL

Hill Top Research, Inc. Main and Mill Streets Miamiville, Ohio 45147

> November 1, 2002 Page 1 of 12



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants Protocol - Rough Plastic

Hill Top Research Confidential

### TEST FACILITY AND INVESTIGATIVE PERSONNEL CON'T. 4.0

Telephone No:

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### 5.0 APPLICABLE REGULATION

Federal Insecticide, Fungicide and Rodenticide Act (40 CFR Part 158) EPA Interim Guidance "Non-Residual Sanitization of Hard Inanimate Food Contact Surfaces Using Pre-Saturated Towelettes (April 12, 2001)"

### 6.0 RESEARCH STANDARDS

This study will be run according to Good Laboratory Practice Standards (40 CFR Part 160). In-Life Phase and Final Report audits will be conducted by the Quality Assurance Unit of Hill Top Research, Inc.

### **EXPERIMENTAL DESIGN** 7.0

Plastic pans identified as "Nalgene® Polypropylene Pan" [pre-sterilized] are inoculated with a specific number of the test bacteria, containing 5% Fetal Bovine Serum to represent a moderate soil load. Plastic pans were selected as they allow the test surface to be thoroughly submerged in the recovery/neutralization media thus achieving reproducible recovery and immediate neutralization. A deeply grooved pattern was mechanically pressed into the bottom of the plastic pans to simulate a rough plastic surface. The inoculum is placed on the rough area (5 and 7/8" X 8") on the bottom of the pan. A sufficient number of pans are inoculated to represent the specified wiped surface area for testing (wiped surface area per pan measures approximately 13.25" X 11" or 1 square foot). The pans are then wiped with a wipe (12" X 12") containing the chemical germicide for a specified period of time. At predetermined exposure time(s), 30 seconds, the remaining chemical agent on the pan is inactivated, and the surviving bacteria are enumerated. The percent reduction in numbers of test bacteria is then calculated. The percent reduction in numbers of bacteria is calculated from a positive control.

November 1, 2002 Page 2 of 12



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants Protocol - Rough Plastic

Hill Top Research Confidential

# 8.0 PROPOSED EXPERIMENTAL STARTING AND EXPERIMENTAL TERMINATION DATES

Proposed Experimental Starting Date:

November 6, 2002

Proposed Experimental Termination Date:

November 8, 2002

Proposed Completion Date:

December 6, 2002

### 9.0 TEST SUBSTANCE IDENTIFICATION

Two lots of the test substance identified as Kimberly Clark Corporation EPA Registration Number 9402-9 will be used for testing: Code 7345-90A (Saturation date 10-8-02) and Code 7345-91A (Saturation date 10-8-02). The lots of test substance were transferred from Hill Top Research Study Number 02-120917-106 and will be assigned Hill Top Research codes for the generation of the test data.

### 10.0 TEST SUBSTANCE CHARACTERIZATION

The sponsor will assume responsibility for test substance characterization according to 40 CFR Part 160.105.

### 11.0 TEST SYSTEM JUSTIFICATION

The test system is designated by federal regulations: EPA Interim Guidance "Non-Residual Sanitization of Hard Inanimate Food Contact Surfaces Using Pre-Saturated Towelettes (April 12, 2001)."

### 12.0 TEST SYSTEM IDENTIFICATION

The test organism to be used in this study will be Shigella boydii, ATCC 9207 with 5% Fetal Bovine Serum incorporated as the soil load according to EPA Draft Method Guidance #02, April 12, 2001. These organisms will be assigned a unique code to provide for the correct generation of data.

### 13.0 TEST PROCEDURE

13.1 The study will be conducted according to the Germicidal and Detergent Sanitizing Action of Disinfectants Method as described in Chapter 6, Disinfectants, Official Methods of Analysis of AOAC International, 17th Edition, 2000 Section 6.3.03 (Appendix I) with modifications and the EPA Interim Guidance "Non-Residual Sanitization of Hard Inanimate Food Contact Surfaces Using Pre-Saturated

November 1, 2002 Page 3 of 12



HTR Ref No.: 02-120918-106 Modified AOAC Germicidal and Detergent Sanitizing Action of Disinfectants Protocol - Rough Plastic Hill Top Research Confidential

### 13.0 TEST PROCEDURE CON'T.

Towelettes (April 12, 2001)." Records will be maintained to verify compliance with these procedures, and any approved modifications to these procedures.

- 13.2 The wipes (test substance) will be tested against the test organisms containing 5% Fetal Bovine Serum to represent a soil load.
- 13.3 The wipes will be tested as received from the sponsor. The wipe will be 12" X 12" and removed from the roll immediately prior to testing. [Two sheets will be removed from the roll and discarded prior to removing the test sheet.] One (12" X 12") wipe will be used to wipe 4 pans/carriers. One wipe will be used to wipe the specified total test surface area represented by wiping consecutive carriers. The surface area wiped per carrier measures approximately 13.25" X 11" allowing each pan to represent 1 square foot of wiped test surface (4 pans represent 4 square feet of surface area wiped). [The wipe will be folded two times in half so that each separate folded portion of the wipe will wipe one of each of the four test pans.]
- 13.4 Exposure conditions will be for 30 seconds at 23  $\pm$ 1°C after a wiping time of 30 seconds.
- 13.5 The neutralizer will be AOAC Neutralizer Blanks with Sea Sand in 400-mL amounts. Neutralizer effectiveness will be determined according to Hill Top Research Standard Operating Procedure 11-DEPP-20-0015A with both test organisms using the plastic pan. The neutralizer will be added to the wiped pan and then the surface will be rubbed [~34 times in the vertical position, ~18 times in the horizontal position, and once around the entire edge (repeat 2 times) in a period of approximately 1 minute] with a sterile rubber policeman to remove the bacteria.
- 13.6 Other modifications to the AOAC method are as follows:
  - The organisms will be harvested using 1.5 mL of AOAC Phosphate Buffer
    Dilution Water per bottle instead of 3.0 mL as listed in Section 6.3.03D of the
    AOAC method.
  - 2) A 0.8-mL aliquot of the adjusted test culture suspension (~1.5±0.5 x 10<sup>8</sup>) will be used to inoculate each gridded test surface (5 and 7/8 "X 8") so that each area of test surface (1 pan/1 square foot) will be inoculated to contain approximately 2.8 X 10<sup>7</sup> CFU/carrier for 1 pan [1 sq. ft.] yielding a count of ~7.5-12.5 X 10<sup>7</sup> CFU/total surface area. The inoculum will be spread evenly over the pan in a drop-wise fashion so that rows of drops are applied to the rough area on the bottom of each plastic pan (a pattern consisting of ~8-9 drops X ~10-11 drops). The pans will then be allowed to air dry for 40 min at 37±2 °C and a relative humidity of at least 50%. The pans will be loosely covered with aluminum foil during the drying process.

November 1, 2002 Page 4 of 12



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants Protocol - Rough Plastic

Hill Top Research Confidential

### 13.0 TEST PROCEDURE CON'T.

3) Numbers controls for each test organism will be conducted using four pans to give the final count per total surface area (4 square feet).

4) Growth will be confirmed by macroscopic examination rather than the method

listed in Section 6.3.03J of the AOAC method.

- 5) The recovery medium will be Tryptone Glucose Extract Agar with 25 mL/L AOAC Stock Neutralizer. Incubation will be at 35 ± 2 °C for 48 ± 2 hours. Plating will be conducted within thirty minutes of neutralizing the test substance by the Pour Plate Method. Two, 10-mL amounts (10<sup>-1</sup>) of the AOAC Neutralizer Blanks with Sea Sand will be plated across three plates and duplicate 1-mL and 0.1-mL amounts (10<sup>-2</sup> and 10<sup>-3</sup> dilutions) will be pour plated. [AOAC Phosphate Buffer Dilution Water with Sea Sand (400-mL) and AOAC Phosphate Buffer Dilution Water (9-mL) and Tryptone Glucose Extract Agar will be used for the numbers controls.] Plate counts will be conducted in duplicate (a + b) and averaged for each pan. Colony counts per milliliter will be multiplied by 4 to yield Colony forming Units (CFU's) per square foot.
- 13.7 Observations of conditions during the test will be recorded in the study records.
- 13.8 Plate counts will be conducted on the expressed fluid from the wipes immediately after wiping the pans (carriers). Dilutions will be conducted in 9.9 mL or 9 mL volumes of AOAC Neutralizer Blanks with plating as outlined in Section 13.6(4 with modifications to account for use of 100 mL of diluent in place of 400 mL of diluent.
- 13.9 The percent reduction in numbers of test bacteria per square foot of surface area (each pan) will be determined as follows:

For each pan

% Reduction/ 1sq. ft. =

[Mean (Avg.) of Numbers Control (Pan 1a + ...Pan 4b) - Survivors of Individual Pan (a + b)]

8 2 X 100

[Mean (Avg.) of Numbers Control (Pan 1a + ... Pan 4b)]

13.10 The percent reduction in numbers of test bacteria per total surface area will be determined as follows:

November 1, 2002 Page 5 of 12



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent
Sanitizing Action of Disinfectants Protocol – Rough Plastic
Hill Top Research Confidential

## 13.0 TEST PROCEDURE CON'T.

For each of the 4 pan set (4 square feet)
% Reduction/ 4 square feet. =

[Sum of Numbers Control (Pan 1a + ... Pan 4b)] - [Sum of Survivors (Pan 1a + ... 4 b)]
2

X 100

[Sum of Numbers Control (Pan la + ... Pan 4b)]

13.11 The percent reduction in numbers of test bacteria per total surface area, including the surviving organisms recovered in the expressed fluid from the wipe, will be determined as follows:

For each of the 4 pan set (4 square feet)
% Reduction/ 4 square feet. =

[Sum of Numbers Control (Pan 1a + ... Pan 4b)] - [(Sum of Survivors Pan 1a + ... 4 b/2) + Survivors from Expressed Fluid]

X 100

[Sum of Numbers Control (Pan 1a + ... Pan 4b)]

## 14.0 STATISTICAL METHOD

No statistical analysis is required to interpret the results of this study.

### 15.0 REPORT

A draft report will be issued, for review by the sponsor, prior to issuing the final report. The report will include (but not be limited to) identification of the test organism, test procedure, protocol modification (if any), identification of the test material, solvent (if any), test concentration, subculture media, results, and summary.

## 16.0 DATA RETENTION

The final report and a copy of the raw data will be sent to the sponsor following completion of the study. All records that would be required to reconstruct the study and demonstrate adherence to the Protocol will be maintained. Following completion of the study, the original raw data and the original of the final report will be maintained indefinitely in the form of hard copy to comply with EPA record keeping regulations. The testing facility will retain a copy of these study records in the form of microfilm.

Upon completion of testing, the test substance will be held for one month and then destroyed; or, at your request and cost, sent back to you.

November 1, 2002 Page 6 of 12



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants Protocol - Rough Plastic

Hill Top Research Confidential

17.0 **NOTICE** 

If it becomes necessary to make changes in the approved protocol, the revisions and reasons for change will be documented, reported to the sponsor and will become part of the permanent file for that study.

Similarly, the sponsor will be notified as soon as is practical whenever an event occurs that is unexpected and may have an effect on the validity of the study.

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HTR Ref No.: 02-120916-106 Modified AOAC Germicidal and Detergent Sanitizing Action of Disinfectants Protocol - Rough Plastic Hill Top Research Confidential PROTOCOL APPROVAL FORM MICROBIOLOGICAL SERVICES DIVISION 18.0 HILL TOP RESEARCH, INC.

REFERENCE CODE PROTOCOL TITLE DISF\PRO\GERM.SAN\KIMC Modified AOAC Germicidal and Detergent Sanitizing Action of Disinfectants-Rough Plastic

PROTOCOL APPROVED FOR: HILL TOP RESEARCH, INC.

BY:

Cathleen A. Baxter, B.S.

1.0/102 Date

Study Director

Microbiological Services Division

Protocol Approved By (Sponsor's Representative):

Signed: Rhonda D. Jones

Title: Agent for Kimberly-Clark Corporation Scientific & Regulatory Consultants, Inc.

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November 1, 2002 Page 8 of 12

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HTR Ref No.: 02-120918-106 Modified AOAC Germicidal and Detergent Sanitizing Action of Disinfectants Protocol – Rough Plastic Hill Top Research Confidential

APPENDIX I

**AOAC Method** 

November 1, 2002 Page 9 of 12



HTR Ref No.: 02-120918-106

Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants Protocol - Rough Plastic

Hill Top Research Confidential

Disinfectants Chapter 6, p. 10

AGAC OFFICIAL METHODS OF ANALYSIS (2000)

### 6.3.03

### **AOAC Official Method 960.09** Germicidal and Detergent Sanitizing Action of Disinfectants First Action 1960 Final Action

(Sultable for determining minimum concentration of chemical that can be permitted for use in sanitizing precleaned, nonporous food contact surfaces. Minimum recommended starting concentration is 2-4x this concentration. Test also determines maximum water hardness for claimed concentrations. As control, check accuracy of hard-water tolerance results with pure C<sub>14</sub> alkyl dimethyl bennyl am-monium chloride at 700 and 900 ppm (µg/mL) hardness, and pure C<sub>14</sub> alkyl dimethyl bennyl ammonium chloride [Cetalkonium Chlo-ride], at 400 and 550 (µg/mL) ppm hardness, expressed as CaCO<sub>2</sub>.)

### A. Respents

(a) Culture media.—(1) Nutrient agar A.—Boil 3 g beof extract, 5 g peptone (from Difco No. 0118 or equivalent; special grades must nothe used), and 15 g salt-free ager in 1 L.H<sub>2</sub>O. Do not use premixed, dehydrated media. Tube, and autoclave 20 min at 121°C. Use for daily transfer of test culture. (2) Nutrient ager 8.—Prepare as above but use 30 g ager. Use for growing test cultures in French square bottles. (3) Nutrient ager (AOAC).—See 955.11A(c) (are 6.1.01). Use for preparing stock culture slants.

(b) Subculture media.—(1) Use tryptone glucose extract agar (Difco No. 0002), adding 25 ml. atock neutralizer, (c)/L. (2) Tryptone glucose extract agar (Difco).

(e) Nestralizer stock solution.—Mix 40 g Lecithin (Alcolec Granules, American Lecithin, PO Box 1908, Danbury, CT 06813. USA [25-50 kg containers only] or Advanced Lecithin Products, PO Box 677, Danbury, CT 06804, USA), 280 mL polysorbate 80, and 1.25 mL phosphate buffer, (e); dilute with H<sub>2</sub>O to I L and adjust to pH 7.2. Dispense in 100 mL portions and autoclave 20 min at 121°C

(d) Neutralizer blanks.—For use with \$200 oom quatemary am rm compound. Mix 100 mL neutralizer stock solution, (c), 25 mL 0.25M phosphate buffer stock solution, (e), and 1675 mL  $\rm H_2O$ Dispense 9 mL portions into  $20 \times 150$  mm tubes. Autoclave 20 min at 121°C.

(e) Phasphate buffer stock solution.—0.25M. Dissolve 34.0 g KH<sub>2</sub>PO<sub>4</sub> in 500 mL,  $H_2$ O, adjust to pH 7.2 with 1M NaOH, and dilute

(I) Phosphate buffer dilution water.-Add 1.25 mL 0.25M phosphate buffer stock solution, (e), to 1 L H<sub>2</sub>O and dispense in 99 mL portions. Autoclave 20 min at 121°C.

(g) Test organisms.-Use Escherichia coli ATCC No. 11229 or Staphylococcus aureus ATCC 6538. Incubate 24 and 48 h, respectively. Maintain stock cultures on nutrient agar (AOAC), (a)(3), at refrigerator temperature.

### B. Resistance to Phenol of Test Cultures

Determine resistance to phenol at least every 3 months by 955.11 (see 6.1.01). Resistance of E. coli should be equivalent to that specified for S. typhi in 955.11D (see 6.1.01) and that for S. aureus equivalent to that specified for this organism in 955.12 (see 6.1.02); also, use procedures under 991.48A(b) (see 6.2.03) for S. aureus.

### C. Apperatus

- (a) Glassware.—250 mL wide-mouth Erlenmeyers; 100 mL graduate; Mohr, serological, and/or bacteriological (APHA specification) pipets; 20 x 150 mm test tubes. Sterllize at 180°C in hot air oven ≥2 h.
- (b) Petri dishas.-Sterile.
- (c) French square bottles.—175 mL, flint glass.
  (d) Water bath.—Controlled at 25°C.

### D. Preparation of Culture Suspension

From stock culture inoculate tube of nutrient agar A, A(a)(I), and make ≥3 consecutive daily transfers (≤30), incubating transfers 20-24 h at 35-37°C. Do not use transfers >30 days. If only I daily transfer has been missed, no special procedures are required; if 2 daily transfers are missed, repeat with 3 daily transfers.

Prepare 175 ml. French square culture bottles containing 20 ml. nutrient agar B, A(a)(2), autoclave 20 min at 121°C, and let solidify with bottle in horizontal position. Inoculate culture bottles by washing growth from slam with 5 mL phosphate buffer dilution H<sub>2</sub>O, A(f), into 99 mL phosphate buffer dilution H<sub>2</sub>O, and adding 2 mL of this suspension to each culture bottle, tilting back and forth to dis-tribute suspension; then drain excess liquid. Incubate 18-24 h at 35-37°C, agar side down. Remove culture from agar surface of 4 or more bottles, using 3 mL phosphate buffer dilution H<sub>2</sub>O and glass beads in each bottle to suspend growth. Filter suspension through Whatman No. 2 paper prewet with 1 mL sterile phosphate buffer, and collect in sterile nube. (To hasten filtration, rub paper gently with sterile policemen.) Standardize suspension to give average of 10 × 10° organisms/mi. by dilution with sterile phosphate buffer dilution H2O, A(f).

Table \$60,09A Percent light transmission at various wavelengths corresponding to bacterial concentrations

	% Light transmission with filters, nm						Average bacterial
370	420	490	530	550	560	650	countril
7.0	4.0	6.0	6.0	6.0	7.0	8.0	13.0 × 10 <sup>9</sup>
6.0	5.0	7.0	7.0	7.0	6.0	9.0	11.5
9.0	6.0	8.0	8.0	8.0	9.0	10.0	10.2
10.0	7,0	9.0	9.0	9.0	11.0	11.0	8.6
11.0	8.0	10.0	10.0	10.0	12.0	13.0	. 7.7
13.0	9.0	12.0	12,0	12.0	13.0	15.0	6.7

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AOAC OFFICIAL METHODS OF ANALYSIS (2000)

DISAFECTANTS Chapter 6, p. 11

Table 960.098 Preparation of Bas04 suspensions corresponding to bacterial concentration

Standard No.	2% BaCl <sub>2</sub> solution, mL	1% H <sub>2</sub> SO <sub>4</sub> (v/v) solution, mL	Average bacteria count/mL
1	4.0	96.0	5.0 × 10 <sup>9</sup>
2	5.0	95.0	7.5
3	6.0	94.0	8.5
4	7.0	93.0	10.0
Б	8.0	92.0	12.0
6	10.0	90.0	13.5
7	12.0	88.0	15.0

If Lumetron colorimeter is used, dilute suspension in sterile Lumetron tube to give % T according to Table 960.09A.

If McFerland nephelometer and BaSO, standards are used, select 7 tubes of same id as that containing test culture suspension. Place 10 mL of each suspension of BaSO, prepared as indicated in Table 960.09B, in each tube and seal tube. Standardize suspension to correspond to No. 4 standard.

### E. Synthetic Hard Water

Prepare Solution I by dissolving 31.74 g MgCl<sub>2</sub> (or equivalent of hydrates) and 73.99 g CaCl<sub>2</sub> in boiled distilled H<sub>2</sub>O and diluting to 1 L. Prepare Solution 2 by dissolving 56.03 g NaHCO<sub>2</sub> in boiled distilled H<sub>2</sub>O and diluting to 1 L. Solution I may be heat scrilized: Solution 2 must be sterilized by filtration. Place required amount Solution I in sterile 1 L. flask and add 2600 mL sterile distilled H<sub>2</sub>O; then add 4 mL Solution 2 and dilute to 1 L with sterile distilled H<sub>2</sub>O. Each mL Solution I will give a water equivalent to ca 100 ppm of hardness calculated as CMCO<sub>3</sub> by formula:

### Total hardness as ppm (µg/mL) CaCO<sub>3</sub> = 2.495 × ppm (µg/mL) Ca + 4.115 × ppm (µg/mL) Mg

pH of all test waters \$2000 ppm (ug/nL) hardness should be 7.6-8.0. Check prepared synthetic waters chemically for hardness at time of tests, using following method or other methods described in APHA, Standard Methods for the Examination of Water and Wattewater 20th Ed., 1992.

### F. Hardness Method

(a) EDTA standard solution.—Dissolve 4.0 g Na<sub>2</sub>H<sub>2</sub>EDTA-2H<sub>2</sub>O and 0.10 g MgCl<sub>2</sub>-6H<sub>3</sub>O in 800 mL H<sub>2</sub>O and adjust by subsequent dilution so that 1 mL of solution is equivalent to 1 mg CaCO<sub>3</sub> when titrated as in (c). Check EDTA solution after preparation or, if commercially purchased, against CaCO<sub>3</sub> standard at least every 2 months.

(b) Calcium standard solution.—1 mL = 1 mg CaCO<sub>2</sub>. Weigh 1.00 g CaCO<sub>3</sub>, died overnight or longer at 105°C, into 500 ml. Eriemmeyer and add dilute HCl through funnel until CaCO<sub>3</sub> is dissolved. Add 200 mL H<sub>2</sub>O<sub>3</sub> boil to expel CO<sub>2</sub>, and cool. Add few drops methyl red indicator and adjust color to intermediate orange with dilute NH<sub>2</sub>OH or HCl as required. Transfer quantitatively to 1 L volumetric flask and dilute to volume.

(c) Determination.—Dilute 5-25 mL test sample (depending on hardness) to 50 mL with H<sub>2</sub>O in Ettenmeyer or casserole. Add 1 mL buffer solution (67.5 g NH<sub>2</sub>Cl and 570 mL NH<sub>2</sub>OH diluted to 1 L with H<sub>2</sub>O), 1 mL inhibitor (5.0 g Nn<sub>2</sub>S-9H<sub>2</sub>O or 3.7 g Nn<sub>2</sub>S-SH<sub>2</sub>O dissolved in 100 mL H<sub>2</sub>O), and one or 2 drops indicator solution (0.5 g Chrome Black T in 100 mL 60-80% sloohed). Tirrate with EDTA standard solution slowly, stirring continuously, until last reddish tinge disappears from solution, adding last few drops at 3-5 s intervals.

# Hardness as mg CaCOy/L \* (mL standard solution × 1000)/mL test sample

### G. Preparation of Test Samples

Use composition declared or determined as guide to test sample weight required for volume sterile  $H_2O$  used to prepare 20 000 ppm ( $\mu g/mL$ ), solution. From this stock dijution, transfer is  $H_1$  into 99 mL of the water to be used in test to give concentration of 200 ppm ( $\mu g/mL$ ). In making transfer, fill 1 mL pipet and drain back into stock solution; then refill, to correct for adsorption on glass. After mixing, discard 1 mL to provide 99 mL of the test water in H.

### H. Operating Technique

Measure 99 mL water to be used in test, containing bactericide at concentration to be tested, into chemically clean, sterile, 230 mL wide-mouth Erlenmeyer and place in constant temperature bath until it reaches 25°C, or≥20 min. Prepare duplicate flasks for each germicide to be tested. Also prepare similar flask containing 99 mL sterile phosphate buffer dilution H<sub>2</sub>O, A(f), as "initial numbers" control.

Add 1 mL culture suspension to each test flask as follows: Whirl flask, stopping just before suspension is added, creating enough residual motion of liquid to prevent pooling of suspension at point of contact with test water. Add suspension midway between center and edge of surface with tip of pipet slightly immersed in test solution. Avoid touching pipet to neck or side of flask during addition. Transfer 1 mL portions of this exposed culture to neutralizer blanks exactly 30 and 60 s after addition of suspension. Mix well immediately

For "numbers control" transfer, add I mL culture suspension to 99 mL sterile phosphate dilution H<sub>2</sub>O in same manner. In case of numbers control, plants need be made only immediately after adding and mixing thoroughly \$30 s. (It is advantageous to use milk pipers to add culture and withdraw test samples.)

Plate from neutralizer lube to agus, using subculture medium A(b)(1) for quaternary ammonium compounds and A(b)(2) with numbers control. Where 0, 1 ml, portions are plasted, use I ml, tipet graduated in 0.1 ml, intervals. For dilutions to give countable plates, use phosphate buffer dilution H<sub>2</sub>O, A(f). For numbers control, use following dilution procedure: Transfer I ml exposed culture (i ml, culture suspension transferred to 99 ml, phosphate buffer dilution H<sub>2</sub>O in H<sub>2</sub>O bath) to 99 ml, phosphate buffer dilution H<sub>2</sub>O, A(f), (dilution I). Shake thoroughly and transfer I ml, dilution I to 99 ml, phosphate buffer dilution H<sub>2</sub>O (dilution 3). Shake thoroughly and transfer I ml, dilution H<sub>2</sub>O (dilution 3). Shake thoroughly and transfer I ml, and four I ml, and four I ml, and four I ml, and four I ml. and four 0.1 ml. aliquots from dilution 3 to individual sterile Petri dishes.

For test samples, use following dilution procedure: Transfer 1 mL exposed culture into 9 mL neutralizer, A(d). Shake and transfer four 1 mL and four 0.1 mL aliquots to individual serile Petri dishes. For numbers control, use subculture medium A(b)(2); for tests with quantum

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> HTR Ref No.: 02-120918-106 Modified AOAC Germicidal and Detergent Sanitizing Action of Disinfectants Protocol - Rough Plastic Hill Top Research Confidential

> > AOAC OFFICIAL METHODS OF ANALYSIS (2000)

Chapter 6, p. 12

ternary summation compounds, use medium A(b)(I). Cool ages solidify, and then invert and incubate 48 h at 35°C before counting. ounds, tate medium A(b)(I). Cool ager to

To be considered valid, results must meet standard effectiveness: 99.999% reduction in count of number of organisms within 30 s. Re-99.9990 remotion to count of manner of organisate water to opport results according to actual count and paycent reduction over manners control. Counts on numbers control for germinide test mixters should fall between 75 and  $125 \times 10^6/mL$  for percent reducti to be considered valid.

(a) Neutralines.--Plate 1 mL from previously unope

(a) Neutrollors.—Plate I mL from previous; unoposes une.

(b) Water.—Plate I mL fram each type of water used.

(c) Sterile distilled water.—Plate I mL After counting plates, confirm that surviving organisms are E. coli by transfer to brillant green bile broth fermentation tubes or lactors both and EMB agar; icm S. aweus by microscopic examination.

es: Am. J. Public Health 38, 1405(1948). L Milk Food Technol. 19, 183(1956). Fed. Regist. 21, 7020(1956). JAOAC 41, \$41(1958); \$6, 308(1973).

**AOAC Official Mathod 961.02** emicidal Spray Products as Disinfectants First Action 1961 Final Action 1964

(Suitable for descripting effectiveness of sprays and pressurist spray producing as spot disinfectants for conteminated surfaces.)

negents specified in 991.47A(s) and (f) Use culture w (see 62.02); 991.48A(a) (see 62.03); and 991.49A(a) and (b) (rec 6.2.05).

Use as best organisms l'incoppieur memagropolisme. No. 9513, prepared as in 955.17D (see 6.302), to which has been added 0.02 mL oxyl-phenoty-poly-shared (Thinto x100, Un-tion Carbide Corp.)/10 mL supprension to fuciliste spreading. Selecion Carbide Corp.)/10 ml. av ion Carbicle Corp.)/10 mL subpension to Facilities speciants, amount of the interneus of ATCE No. 10708, maintained as in 991.474,00 (cee 6.202), Susphylacoccus authors ATCE No. 6538, maintained as in 991.484,00 (cee 6.202), and Pacuslamonas aeruginoss ATCE No. 15442, maintained as in 991.484(c) (cee 6.2.05), incubate all bacterials. tacial cultures for 48 b, except porm

Use apparates specified in 991.678(b), (b), (c), (a), and (c) (see 6.2.02), and in addition:

(a) Capillary pipets.—0.1 ml., gradual Sterilize in air oven 2 h at 180°C

corresive, 25 x 25 mm (1 × 1 in.). (b) Microscope slides.—No or 18 × 36 cnm glass slide. Sterilize by placing individual slides in Petri dish nasted with 2 pieces 9 cm Oline paper (Whatman No. 2, or equivalent) in air oven 2 h at 180°C.

(c) Bacteriological culture tubes.—Pyrex, 32 x 200,000 (Belico Hass, Inc., PO Box B, Vineland, NJ 08360, USA).

(d) Metal forceps.—Sharp points, straight, 115 mm

Thoroughly shake 48 h nutrient broth cultures of tholoressuir and 5. sureus and let settle 10 min. For 5. Photerassuir and 5. surems and let sattle 10 min. For P. arrughosa, follow preparation of culture under 991.43A(c) (see £,2.05). With sterile capillary pipet or sterile 4.0 mm loop, transife 0.0 m la culture soot 1 as in sterile test silde in Petri dish and imboediately spread uniformly over entire urea. Cover dish immediately and repeat operation until 12 slides have been prepared for pach organism. (Use 2 slides as control.) Dry all slides 30-40 min at 37°C. pared for each organ 30-40 min at 37°C.

30–40 min at 37°C.

Spray [Onlides for specified time and distance. If no time or distance specified, use 10 s at 1. ft. (30 cm). Hold each slide 10 min, draid off access liquid, and transfer slide to individual 32 × 200 mm hybe containing 20 mL appropriate subcolume medium, 935.11A(s) (see 6.1.01), with fiamed forceps. Shake culture thoroughly If broth appears cloudy after 30 min, make subculture to fresh individual tubes of subculture broth. Transfer 2 unapprayed slides, as vishility controls, to individual subculture babes in same manker.

Incubate all tubes used for primary and secondary transfers 46 h at 37°C. Read as 4 (growth) or (no growth). Killing of test organisms in 10 of 10 trials is produmptive evidence of disinfecting action.

For procedures to be followed in assuring standard colluves, for 5. cholernessuit, see 394.47A(b) (see 6.2.02), for 5. aurens, 591.48A(b) (see 6.2.03) for P. aeraginosa, see 391.48A(c)

S. choloroscuis, see 394.47A(b) (see 6.2.02), for S. aureur, 391.48A(b) (see 6.2.03) for P. auruginosa, see 991.49A(c) (see 6.2.05). For T. auruginosa, see 955.17A and D (see 6.2.02). If there is reason to believe that seck of growth in subtransfer tubes is due to bacteriostasia inoculate all insubsted subculture tubes with loop assed inoculation of respective best culture and resinculates. Growth of times inducia eliminates besteriostasia sa cases of lack of growth. If there is question as to possibility of contamination as source of growth in subculture tubes, make gram stains and/or subculture for identification, according to respective test culture. tive test culture.

spective test cantre.

If fungicidal activity as well as geomicidal activity is involved,
use test suspension of T. menter/tophyses spores, 955.17D
(see 6.3.02), and propers 12 slides, using 0.01 mL standard spore
suspension, spraying and subculturing exactly as above. Make
subcultures in glucose broth, 955.178 (see 6.3.02), incubating 7 days at 25-30°C.

Soap Chem. Spec. 38(2), 69(1967).

ACAC Official Method 966. Sportcidal Activity of Disinfect

First Action 1986 Final Action 1987

(Suitable for determining spotleidal activity of it gasoous chamiculs. Applicable to germicides for de pressure or absence of spotleidal activity against spore-forming bacteria in various situations and pots eacy as sterilizing agent.)

(a) Culture media....(1) Soil extract natrient broth 1 ib (454 g) garden soil in 1 L H<sub>2</sub>O, filter several times the No. 588 paper, and dilute to volume (pH should be ≥5.2). Add\ g

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HILL TOP RESEARCH, INC.

MAIN AND MILL STREETS **MIAMIVILLE, OHIO 45147** 

PROTOCOL AMENDMENT #1 Modified AOAC Germicidal and Detergent

Sanitizing Action of Disinfectants - Rough Plastic

One Step Cleaner Sanitizer

HTR STUDY NO.:

02-120918-106

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### PROTOCOL AMENDMENT:

1. At the request of the sponsor representative on November 5, 2002, Section 13.11 of the protocol will be amended. The current section will be deleted and replaced with the following statement:

13.11 The percent reduction in numbers of test bacteria surviving in the expressed fluid from the wipe will be determined as follows:

[Sum of Numbers Control (Pan la + ... Pan 4b)] - [Survivors from Expressed Fluid] X 100

[Sum of Numbers Control (Pan 1a + ... Pan 4b)]

2. This change is being made to follow the May 22, 2002 Efficacy Review from USEPA requesting that the percent reduction in numbers of bacteria surviving in the expressed fluid from the wipe be calculated for EPA Registration Number 9402-O.

APPROVED FOR: HILL TOP RESEARCH, INC.

BY: athleen A. Baxter, B.S.

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COPY

ACCEPTED BY: Kimberly Clark Corporation

BV.

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Page 2 of 2 02-120918-106